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LETTER

Triose phosphate limitation in photosynthesis models reduces leaf photosynthesis and global terrestrial carbon storage

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Abstract

Triose phosphate utilization (TPU)-limited photosynthesis occurs when carbon export from the Calvin-Benson cycle cannot keep pace with carbon inputs and processing. This condition is poorly constrained by observations but may become an increasingly important driver of global carbon cycling under future climate scenarios. However, the consequences of including or omitting TPU limitation in models have seldom been quantified. Here, we assess the impact of changing the representation of TPU limitation on leaf- and global-scale processes. At the leaf scale, TPU limits photosynthesis at cold temperatures, high CO₂ concentrations, and high light levels. Consistent with leaf-scale results, global simulations using the Community Land Model version 4.5 illustrate that the standard representation of TPU limits carbon gain under present day and future conditions, most consistently at high latitudes. If the assumed TPU limitation is doubled, further restricting photosynthesis, terrestrial ecosystem carbon pools are reduced by 9 Pg by 2100 under a business-as-usual scenario. The impact of TPU limitation on global terrestrial carbon gain suggests that CO₂ concentrations may increase more than expected if models omit TPU limitation, and highlights the need to better understand when TPU limitation is important, including variation among different plant types and acclimation to temperature and CO₂.

Introduction

Photosynthesis is the largest gross carbon flux (~120 Pg C yr⁻¹) between Earth's surface and the atmosphere, enabling terrestrial sequestration of approximately one-third of anthropogenic carbon dioxide (CO₂) emissions (Le Quéré *et al* 2016). Models of leaf-level photosynthesis (e.g. Farquhar *et al* 1980) provide the foundation for estimating global CO₂ uptake by plants, yet there is considerable uncertainty associated with the structure and parameterization of

photosynthesis in these models (Arora *et al* 2013, Bonan *et al* 2012, Rogers 2014, Rogers *et al* 2017a). The recent CMIP5 ensemble of Earth system model (ESM) simulations showcases this uncertainty, with present day (1986–2005) estimates of Northern Hemisphere gross primary production (GPP) ranging from 6–16 Pg C in July (Anav *et al* 2013) and estimates of global cumulative land uptake between 1850 and 2005 spanning ~150 Pg C across all models (Ciais *et al* 2013).

One source of uncertainty in modeled estimates of carbon fluxes arises from differences in how



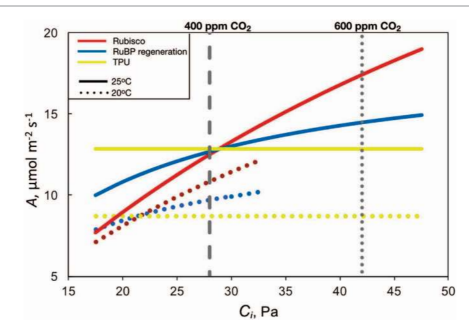


Figure 1. Examples of assimilation versus internal leaf CO $_2$ (A/C_i) curves assuming Rubisco (red), RuBP regeneration (blue) or triose phosphate utilization (TPU; yellow) limitation at either 25° C (solid lines) or 20° C (dotted lines). The vertical dashed line corresponds to 400 ppm ambient CO $_2$ (C_a) and the vertical dotted line corresponds to 600 ppm, assuming that C_i is 0.7 times C_a . The data were generated using the equations and temperature sensitivities in Sharkey (2016), with the maximum rate of carboxylation ($V_{\rm cmax}$) set to 50 μ mol m⁻² s⁻¹, electron transport (J) set to 80 μ mol m⁻² s⁻¹, and TPU set to $^{1}/_{12}$ $V_{\rm cmax}$, the same as the high TPU limitation scenario used in the leaf photosynthesis model and the CLM4.5-BGC simulations. At 25° C and 400 ppm CO $_2$, all three parameters are equally limiting, resulting in no excess capacity. However, at 20° C and 400 ppm CO $_2$, TPU is the most limiting parameter, resulting in unused capacity in Rubisco and RuBP regeneration. At 600 ppm CO $_2$, TPU is most limiting and sets the maximum rate of net photosynthesis (A_n) at both leaf temperatures.

models represent specific plant processes (e.g. Lovenduski and Bonan 2017, Rogers et al 2017a). For example, Rogers et al (2017a) evaluated how representation of C₃ photosynthesis at leaf and canopy scales varied across several global models and included recommendations for future research that may help to reduce model structural uncertainty and improve carbon cycle predictions. Here, we quantitatively assess one source of variation identified by Rogers et al (2017a)—structural uncertainty associated with the inclusion of triose phosphate utilization (TPU) limitation of photosynthesis (described below), which is only represented in a few Earth system models. In particular, we investigate the impact of TPU limitation on CO_2 assimilation at the leaf level and at the global scale for present-day and projected future carbon storage using the Community Land Model (CLM).

Typically, several physiological limitations to C_3 photosynthesis are represented in photosynthesis models, including biophysical constraints (e.g. stomatal and mesophyll conductance) that occur when CO_2 diffuses into and through the leaf, as well as biochemical constraints that occur during CO_2 fixation and utilization (Rogers *et al* 2017a). Two biochemical constraints that are commonly included in models are photosynthetic limitation by carboxylation and electron transport (see figure 1), which are calculated at the leaf-level using equations originally derived by Farquhar *et al* (1980). Carboxylation-limited photosynthesis (A_c) is often limiting under

low CO_2 partial pressure and is a function of V_{cmax} , which is the maximum carboxylation rate of ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) assuming a saturating supply of ribulose 1,5-bisphosphate (RuBP). Electron transport-limited photosynthesis (A_j) describes photosynthesis limited by electron transport that supports the regeneration of RuBP and can also include limitations within the Calvin-Benson cycle other than Rubisco. It is often limiting under higher CO_2 partial pressures, but can be limiting over a broad range of CO_2 at low light (Sharkey 2016). Net photosynthesis (A_n ; μ mol CO_2 m⁻² s⁻¹) is subsequently calculated as the minimum of A_c and A_j :

$$A_n = \min\{A_C, A_i\} - R_d \tag{1}$$

where R_d is non-photorespiratory respiration in the light,

$$A_C = \frac{V_{\text{cmax}}(C_i - \Gamma^*)}{C_i + K_C \left(1 + \frac{o_i}{K_o}\right)}$$
(2)

and

$$A_j = \frac{J(C_i - \Gamma^*)}{4C_i + 8\Gamma^*}. (3)$$

In these equations, $J(\mu \text{mol electrons m}^{-2} \text{ s}^{-1})$ is based on the maximum potential electron transport rate (J_{max}) , C_i is the intracellular concentration of CO₂ (Pa), K_c and K_o are the effective Michaelis-Menten



constants for CO_2 and O_2 , O_i is the O_2 partial pressure (Pa), and Γ^* (Pa) is the CO_2 compensation point in the absence of mitochondrial respiration (Farquhar *et al* 1980).

When both carboxylation and electron transport rates are fast, a third biochemical constraint can limit the rate of photosynthesis. This occurs when carbon is not exported from the Calvin-Benson cycle as quickly as it is fixed and is known as TPU limitation, and TPU limited A is denoted (A_p) . TPU limitation is typically assumed to be negligible if it is included in photosynthesis models and is therefore often omitted. However, TPU limitation may become more prevalent under future climate conditions. When the TPU limitation is included with the biochemical limitations discussed above, A_n can be calculated as:

$$A_n = \min\{A_C, A_i, A_p\} - R_d. \tag{4}$$

A detailed representation of A_p accounts for inefficiencies in carbon recycling during photorespiration:

$$A_p = \frac{(C_i - \Gamma^*) \times 3\text{TPU}}{C_i - (1 + 3 \times a_g) \times \Gamma^*}$$
 (5)

where a_g is a unitless scalar from 0–1 based on gly-colate recycling (with 1 being 100% recycling) within the chloroplast (Ellsworth *et al* 2015, Busch and Sage 2016, von Caemmerer 2000). The metabolic basis of equation (5) is thought to apply in many cases and describes the TPU limitation quite well (Harley and Sharkey 1991, Collatz *et al* 1991, Sellers *et al* 1996, Busch *et al* 2018). When glycolate recycling (a_g) is assumed to be 0, equation (5) simplifies to:

$$A_p = 3\text{TPU}. \tag{6}$$

This simpler version of the calculation for A_p is used in some global models, such as the CLM (see Oleson *et al* 2013) and the Joint UK Land Environment Simulator (JULES; Mercado *et al* 2007). In these equations,

$$TPU = 0.167V_{cmax} \tag{7}$$

so that by using equation (7) to calculate equation (6), $A_p = 0.5 V_{\rm cmax}$. The model formulation from which equation (7) is derived originates from work by Collatz et al (1991) and has been used in some land surface models for many years, even though no empirical data explicitly links TPU to $V_{\rm cmax}$. Using observations available in Wullschleger (1993) for $16 \, {\rm C_3}$ plant species, we calculated the ratio of TPU to $V_{\rm cmax}$ to determine that TPU = $0.132(\pm 0.038)^* V_{\rm cmax}$, suggesting that the current calculation of TPU in models falls on the high end of the observed range, and is perhaps a conservative estimate.

Leaf- and global-scale photosynthetic models typically specify values of $V_{\rm cmax}$ used in equation (2) and sometimes $J_{\rm max}$, which is used to calculate J in equation (3). Because these parameters are the foundation for photosynthesis models, it is important that the values are representative of the vegetation they

are simulating (Rogers 2014). For example, changes in the representation of $V_{\rm cmax}$ within the canopy of an Earth system model can cause estimates of global GPP to change as much as $30\,{\rm Pg}\,{\rm C}\,{\rm yr}^{-1}$ (Bonan *et al* 2011). Field-based estimates of $V_{\rm cmax}$, $J_{\rm max}$, and TPU are frequently derived from measured photosynthetic ${\rm CO}_2$ response curves (A/C_i curves), and the values used in many global-scale models are averages across species using data available from sources like the TRY database (Kattge *et al* 2011). However, TPU is not always included in curve fitting analyses. Nonetheless, studies using non-standard methods find that TPU limitation occurs frequently (Ellsworth *et al* 2015, Yang *et al* 2016).

Available research suggests that TPU limitation is most likely to occur under high CO₂ concentrations, high light, and/or low air temperature relative to growth temperature (Harley and Sharkey 1991, Busch and Sage 2016, Bernacchi et al 2009). Despite this basic understanding, the individual and interactive effects of environmental controls on TPU remain unknown for most species (Busch and Sage 2016), so model representations of TPU-limited photosynthesis are either necessarily over parameterized, represented nonmechanistically, or excluded (Gu et al 2010, Sharkey 2016, Maayar et al 2006, Rogers et al 2017a). The sensitivity of photosynthesis to TPU limitation has rarely been examined across global scales even though it could become an important limitation to carbon uptake under future environmental conditions.

Here, we explore the impact of TPU limitation under various climate conditions by quantifying the effect of including TPU limitation on photosynthesis across multiple scales. Because global scale models depend on leaf-scale photosynthesis calculations, we first use a leaf-scale photosynthesis model to highlight conditions under which modeled TPU would be expected to limit photosynthesis. We then scale the impact of including TPU-limited photosynthesis on global carbon uptake using the full global land model. We conclude by discussing research priorities for improving our understanding of TPU-limited photosynthesis using targeted field campaigns that will help to evaluate and constrain the representation of TPU in photosynthesis models and improve carbon cycle simulations in ESMs.

Methods

We used a global-scale land model and its embedded leaf-scale photosynthesis submodel to explore the sensitivity of A_n and ecosystem carbon storage to TPU limitation under changing environmental conditions. The global model used here is CLM4.5-BGC (Oleson *et al* 2013). Both the leaf submodel and the CLM4.5-BGC calculate TPU-limited A_n (i.e. A_p) using equation (4) above, and include export-limited photosynthesis based on TPU limitation as in equations (6) and



(7) above (Harley and Sharkey 1991). It should be noted that CLM4.5 solves for A_n (equation 4) as a rate colimited by Rubisco-, light-, and export-limitation (Collatz et al 1991, 1992, Bonan et al 2011) to account for variability in photosynthetic limitations across the canopy, which reduces A_n more than any single limitation (Bonan et al 2011) and may have a greater impact at low temperatures. Both the CLM4.5-BGC and the leaf submodel included photosynthetic temperature acclimation based on Kattge and Knorr (2007), with TPU acclimating in the same manner as $V_{\rm cmax}$ (Lombardozzi et al 2015). In CLM4.5-BGC, photosynthetic temperature acclimation occurs between leaf growth temperatures of 11°C-35°C, though the leaf submodel did not impose temperature restrictions on acclimation. The leaf submodel additionally assumed a V_{cmax} at 25 °C ($V_{\text{cmax}25}$) of 50 μ mol m⁻² s⁻¹, Rubisco kinetics from Bernacchi et al (2001), and an intracellular to extracellular CO_2 ratio (C_i/C_a) of 0.7.

We tested the sensitivity of leaf and global photosynthesis to TPU limitation by testing three different TPU limitation scenarios. The unmodified TPU used in CLM4.5, which we term here 'standard TPU limitation,' calculated TPU as one-sixth of $V_{\rm cmax}$ (equation 7) and the export-limited rate of photosynthesis as three times TPU (equation 6, leading to an exportlimited rate = 0.5 V_{cmax} ; Collatz et al 1991). To test a scenario where plants are more limited by TPU limitation ('high TPU limitation'), we modified TPU to be one-twelfth of $V_{\rm cmax}$ (e.g. equation 7 is updated to TPU = 0.0835 $V_{\rm cmax}$, halving the TPU and doubling the TPU limitation by making the export-limited rate in equation $6 = 0.25 V_{\text{cmax}}$). The high TPU limitation scenario brackets the range of observations (TPU = (0.132 ± 0.038) *V_{cmax}; Wullschleger *et al* 1993) and is close to the low-end estimate of this range $(TPU = 0.094*V_{cmax})$. This simulation was anticipated to reduce carbon gain because TPU will more strongly limit photosynthesis. Finally, to test a scenario where plants are less limited by TPU, termed 'low TPU limitation', we modified TPU to be onethird of the $V_{\rm cmax}$ rate (e.g. equation 7 is updated to TPU = $0.334V_{\text{cmax}}$, doubling the TPU so that the export-limited rate in equation $6 = V_{\text{cmax}}$), lessening the importance of TPU and reducing export-limited photosynthesis. The low TPU limitation scenario illustrates the impact of having very weak to no TPU limitation compared to the standard representation used in CLM.

Using the leaf photosynthesis model, we ran simulations with the high and standard representations of TPU limitation at combinations of varying C_a (1–2000 μ mol mol⁻¹), photosynthetically active radiation (PAR; 50–2000 μ mol m⁻² s⁻¹), and leaf temperature (5 °C–35 °C). In our testing, the low TPU simulation that is used in the global simulations does not limit photosynthesis below a C_a value of 2000 μ mol mol⁻¹ (data not shown) and is therefore not discussed in our leaf photosynthesis analysis. From these simulations,

we identified the C_a value at which modeled photosynthesis became TPU-limited under different light and leaf temperature conditions for 'standard TPU' and 'high TPU' limitations.

To test the implications of TPU limitation for the global carbon cycle, we ran the CLM4.5 with active biogeochemistry (BGC; Oleson et al 2013) at 2 degree resolution from 1850 through 2100 using atmospheric forcing data archived from a Community Earth System Model (CESM) simulation that used a historical forcing scenario through 2005 and the representative concentration pathway 8.5 (RCP8.5) forcing scenario through 2100 (Meehl et al 2012). Each of the TPU scenarios described above (standard, low, and high) was tested in a separate simulation. After the modifications were included to adjust TPU, each simulation was spun up using 1850 initial conditions until carbon pools stabilized. Analysis focused on the changes in GPP, as well as the integrated response of total ecosystem carbon, which sums total soil and vegetation carbon pools, to better understand the consequences of the accumulated changes in productivity on terrestrial carbon storage.

Results and discussion

Leaf-level photosynthesis

The results from the leaf model illustrate that the C_a value where leaves become TPU-limited increases with temperature and decreases with PAR (figure 2). TPU limitation occurs across a range of C_a values, from as low as 200 μ mol mol⁻¹ at a leaf temperature of 5 °C, to over 2000 μ mol mol⁻¹ under high temperature and low light conditions (figure 2). Not surprisingly, assuming lower rates of TPU (i.e. high TPU limitation) limited photosynthesis more across the range of environmental conditions tested here, but the patterns remained the same (figure 2(b)) compared to 2(a)).

Overall, this leaf-scale exercise suggests that Earth system models (ESMs) that include TPU limitation in photosynthesis calculations would predict large geographic areas where photosynthesis is limited by carbohydrate export, particularly cold regions during times of high radiation. It is important to note that the leaf submodel used here assumes a static $V_{\rm cmax25}$ value, whereas observations indicate that $V_{\text{cmax}25}$ varies with environmental conditions such as temperature (Ali et al 2015), nitrogen in leaves (Walker et al 2014, Kattge et al 2009), water availability (Reich et al 2007, Ali et al 2015), day length (Bauerle et al 2012), or ozone exposure (Lombardozzi et al 2012). Additionally, the ratio of TPU to $V_{\text{cmax}25}$ can be decreased by factors such as water stress (Vassey and Sharkey 1989, Lawlor 2002), changes in leaf respiration capacity (Dahal et al 2014), and increased leaf nitrogen (Sage et al 1990).

We assumed that the temperature sensitivity of TPU is the same as that of $V_{\rm cmax}$. Although evidence

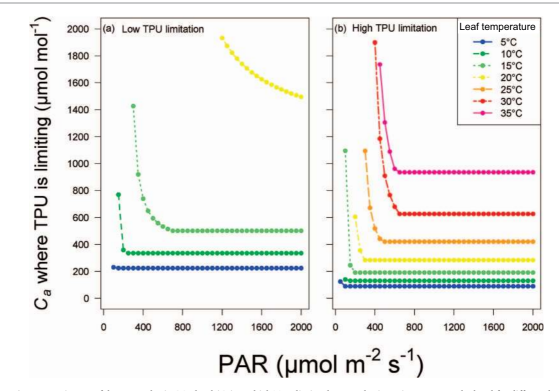


Figure 2. Estimates of the atmospheric CO_2 level (C_a) at which TPU limits photosynthesis. Estimates were calculated for different leaf temperatures (lines), incoming photosynthetically active radiation (PAR) (x-axis), and for (a) low TPU limitation and (b) high TPU limitation using a derivation of the Farquhar model of leaf photosynthesis (Farquhar $et\ al\ 1980$) by Collatz $et\ al\ (1991)$ and modified to include TPU-limited photosynthesis from Harley and Sharkey (1991), similar to equations (5–7) in the text. In all cases, glycolate recycling (a_g) was assumed to be 0. In the low TPU limitation case (a), the model assumed that TPU was one sixth of the maximum rate of carboxylation (V_{cmax}). In the high TPU limitation case (b), the model assumed that TPU was one twelfth of V_{cmax} vas calculated from a rate at 25 °C of 50 μ mol m⁻² s⁻¹ and an Arrhenius temperature response from Kattge and Knorr (2007). J_{max} was calculated from a rate at 25 °C, based on an acclimated ratio of J_{max25} to V_{cmax25} and an Arrhenius temperature response from Kattge and Knorr (2007). The model assumed Rubisco kinetics from Bernacchi $et\ al\ (2001)$ and an intracellular to extracellular CO $_2$ ratio (C_i/C_a) of 0.7.

suggests that physiological processes in plants may acclimate (Ainsworth and Rogers 2007, Sage et al 1989) or even adapt (Watson-Lazowski et al 2016) to elevated CO2 and may therefore change photosynthetic limitations in the future, the leaf model used here, and most Earth System models, do not account for these processes. The introduction of prognostic $V_{\rm cmax}$ (e.g. Xu et al 2012, Ali et al 2016), as planned for CLM5, may alleviate these limitations. Nonetheless, the occurrence of simulated TPU-limited photosynthesis should increase as C_a levels rise, though increasing leaf temperatures could offset this increase. Thus, it is important to improve our understanding of how photosynthesis acclimates to the combination of CO₂ and temperature changes expected in the future.

Global photosynthesis and carbon cycle

Compared to the standard (unmodified) simulation, reducing TPU limitation increased simulated GPP slightly by 2100 under low TPU limitation compared to the standard TPU limitation (+1 Pg C, \sim 2%), whereas high TPU limitation more clearly decreased simulated GPP by 2100 (-3 Pg C, \sim 6%; figures 3(c) and (d)). Similarly, low TPU limitation increased simulated total ecosystem C gain (+4.5 Pg, \sim 4%) by 2100, while

high TPU limitation reduced simulated total ecosystem C gain $(-9.0 \text{ Pg}, \sim 9\%; \text{ figures } 3(a) \text{ and } (b))$. The changes in total ecosystem carbon reflect the integrated C fluxes from changes in productivity.

In CLM4.5 simulations, TPU limits GPP (and the accumulated ecosystem carbon storage, see SI figure 1 available at stacks.iop.org/ERL/13/074025/mmedia) most in the high latitudes (figure 4), particularly in 2100, which is consistent with the results from the leaf photosynthesis submodel that suggests TPU limits photosynthesis more frequently at low temperatures and higher CO₂ (figures 1 and 2). The CLM4.5 simulations also illustrate that high limitation of TPU suppresses GPP by 2000, and increasingly more by 2100. Rogers et al (2017b) showed that $V_{\rm cmax}$ in Arctic vegetation was markedly higher than the values used in current ESMs and approximately double the value used here. Because of the assumed relationship between $V_{\rm cmax}$ and TPU, higher Arctic $V_{\rm cmax}$ might lead to TPU limitation at higher (CO₂) than we found here, resulting in higher GPP and carbon storage. It is important to note that figure 4 shows the change of a change $(\Delta \Delta)$ for the low (figures 4(a) and (c)) and high (figures 4(*b*) and (*d*)) TPU limited scenarios. Thus, differences plotted in figure 4 are a result of both the temporal change in TPU limitation (2100 or 2000-1850) as



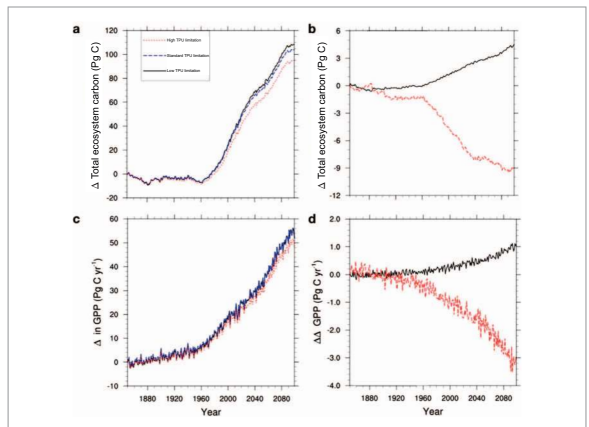


Figure 3. Effects of TPU on (*a*) the change in global total ecosystem carbon relative to 1850, and (*c*) the change in gross primary productivity (GPP) relative to 1850 for the standard representation of TPU in CLM4.5-BGC (dashed blue line), and for simulations with high TPU limitation (short dashed red line) and low TPU limitation (solid black line). The difference in (*b*) global total ecosystem carbon and (*d*) gross primary productivity relative to the standard TPU limitation (as plotted in panels (*a*) and (*c*)) are plotted for the high TPU (short dashed red line) and the low TPU (solid black line) to highlight the magnitude of divergence of each modification.

well as the change in TPU scenario (high or low TPU limitation—standard TPU limitation). Though the spatial patterns of change are similar in the lowand high-TPU limited scenarios, the magnitude of the decrease in GPP (and ecosystem carbon, SI figure 1) is much larger in many regions with high TPU limitation (e.g. $> 100\,\mathrm{g\,C\,m^{-2}}$ yr⁻¹) than the increase in GPP due to reducing TPU limitation. The increased GPP in the low TPU simulation illustrate that the standard parameterization used in CLM4.5-BGC is TPU limited in high latitudes by the end of the twenty-first century.

Regional responses of photosynthesis to changes in TPU limitation may be partly affected by the representation of TPU acclimation to short-term (10-day) changes in mean temperature in CLM (Lombardozzi et al 2015). Though few data are available to verify whether TPU acclimates, TPU is thought to be temperature dependent, and is perhaps more sensitive to temperature changes than V_{cmax} (Yang et al 2016), even though it is often modeled without a temperature function. Temperature is an important determinant of TPU limitation (figure 2), so understanding whether TPU acclimates to temperature changes remains a key research priority. Limitation of photosynthesis under the standard TPU representation in CLM4.5 suggests that TPU limitation likely occurs in models that represent it, especially if acclimation to temperature is not accounted for

The decrease in global (figure 3) and high latitude (figure 4) GPP and ecosystem carbon between the standard and high TPU limitation simulations compared to the low TPU limitation are more intuitive than the relative increases in GPP and ecosystem carbon that occur in tropical regions and a few other locations like Central Europe (figure 4, SI figure 1). In fact, a time-series of GPP and ecosystem carbon in the Amazon Basin illustrates that the absolute values of GPP and ecosystem carbon were lower in simulations that included standard or high TPU limitation throughout the duration of the simulation compared to the low TPU limitation simulation (figure 5). Therefore, the relative increases in response to high TPU limitation (figure 4) result from a larger net carbon gain between 1850 and 2100. It is possible that the maximum GPP and ecosystem carbon gain in tropical regions, for example, is limited by another ecosystem driver like nitrogen availability (see Lombardozzi et al 2015).

Conclusions and future research priorities

Overall, our work illustrates that model estimates of net photosynthesis in leaf- and global-scale models are likely limited by TPU under some environmental



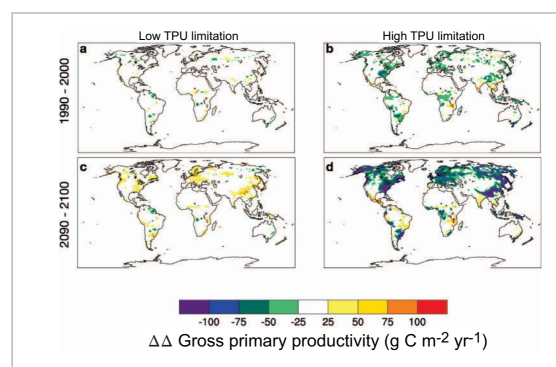


Figure 4. The change in gross primary productivity averaged from 1990–2000 (a) and (b) and 2090–2100 (c) and (d) relative to the 1850–1860 average for CLM4.5-BGC simulations with low TPU limitation (a) and (c) and high TPU limitation (b) and (d) compared to the same change in the simulation with the standard TPU limitation used in CLM4.5. $\Delta\Delta$ notation used here denotes the difference in 1990–2000 or 2090–2100 relative to 1850–1860 (Δ), and the difference between the simulation with either low (a) and (c) or high (b) and (d) TPU limitation relative to the same change in the simulation with standard (unmodified) TPU limitation (Δ). Differences plotted here are therefore due to the temporal change in TPU limitation (2100—1850) as well as to the scenario (high or low TPU limitation).

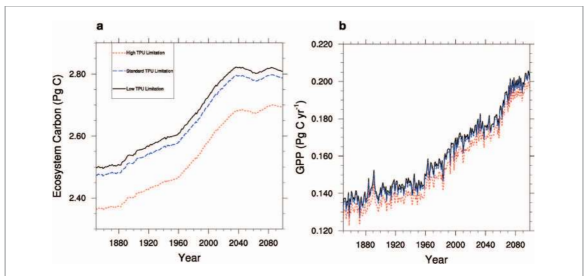


Figure 5. Time series of ecosystem carbon pools (a) and gross primary productivity (GPP) (b) in the Amazon Basin for CLM4.5-BGC simulations with high (red), standard (blue), or low (black) TPU limitation.

conditions (figures 2 and 3), with TPU limitation having the largest impact under high CO_2 in cold regions (figure 4), similar to current theoretical understanding and available observations. Using a global-scale model, we add to current understanding of TPU limitation by identifying times when and regions where TPU limitation may be important. As represented in CLM4.5, the present-day carbon cycle is currently limited by TPU, and the limitation increases under the future RCP8.5 climate scenario (figures 3 and 4). Given

the few available data on TPU, it is difficult to assess the realism of the TPU limitation exerted within the models we tested. The calculation of TPU we use equation (7) is within the range of measured estimates ((0.132 \pm 0.038)* $V_{\rm cmax}$; Wullschleger 1993) and may be a conservative estimate. The observational estimates suggest that TPU limitation is more likely to fall between the standard and high TPU limitation scenarios, so even the ESMs that currently include TPU limitation using the standard formulation may



underestimate its impact. If TPU limitation is underestimated or omitted from photosynthesis calculations in ESMs, then projections of future CO₂ concentration increases are likely also underestimated.

Currently, we lack a robust theoretical understanding of the drivers of TPU limitation under different environmental conditions and over large spatial and temporal scales. Estimates of TPU limitation from in situ measurements may help to constrain carbon cycle uncertainty in model simulations, and measurements collected under environmental manipulations, such as elevated CO2 experiments, can help to better understand whether model projections of future changes in TPU limitation are at all realistic. Our modeling results identify high latitudes as regions where TPU limitation is most likely to occur in the future, and these regions should have the highest priority for measurement campaigns. We recognize that measuring TPU limitation is challenging, but the advent of new techniques, such as combining gas exchange and fluorescence measurements of leaf O2 and CO₂ sensitivities (Busch and Sage 2016, Busch et al 2018) may help with collecting more accurate estimates of TPU, as well as other photosynthetic parameters.

Measurement campaigns should also consider the potential for acclimation of export-limited photosynthesis to environmental conditions thought to be particularly susceptible to high TPU limitation, such as high CO₂, high light, and low temperatures. For example, the temperature sensitivity of TPU documented by Yang et al (2016) suggests that TPU is more sensitive to temperature than $V_{\rm cmax}$. Additionally, recent work using the coordination theory of photosynthesis (e.g. Wang et al 2017, Quebbeman and Ramirez 2016) suggests that plants adjust carboxylation- and electron transport-limited photosynthetic processes such that neither is strongly limiting at current growth conditions. It is possible that similar acclimation of export-limited photosynthesis occurs, though more measurements are needed to fully assess this response. Gaining a better process understanding of TPU limitation through the targeted measurements recommended above is necessary to improve the representation of TPU limitation in photosynthesis models. Ultimately, improved process understanding of TPU limitation may help to constrain the uncertainty in carbon cycle projections in the future, when TPU limitation is expected to more frequently limit photosynthesis.

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References

Ainsworth E A and Rogers A 2007 The response of photosynthesis and stomatal conductance to rising (CO₂): mechanisms and environmental interactions *Plant Cell Environ*. 30 258–70

Ali A A *et al* 2016 A global scale mechanistic model of photosynthetic capacity (LUNA V1.0) *Geosci. Model Dev.* 9 587–606

Ali A A *et al* 2015 Global-scale environmental control of plant photosynthetic capacity *Ecol. Appl.* **25** 2349–65

Anav A, Friedlingstein P, Kidston M, Bopp L, Ciais P, Cox P, Jones C, Jung M, Myneni R and Zhu Z 2013 Evaluating the land and ocean components of the global carbon cycle in the CMIP5 Earth system models *J. Clim.* 26 6801–43

Arora V K *et al* 2013 Carbon–concentration and carbon–climate feedbacks in CMIP5 Earth system models *J. Clim.* **26** 5289–314

Bauerle W L, Oren R, Way D A, Qian S S, Stoy P C, Thornton P E, Bowden J D, Hoffman F M and Reynolds R F 2012
Photoperiodic regulation of the seasonal pattern of photosynthetic capacity and the implications for carbon cycling *Proc. Natl Acad. Sci. USA* 109 8612–7

Bernacchi C J, Rosenthal D M, Pimentel C, Long S P and Farquhar G D 2009 Modeling the temperature dependence of C₃ photosynthesis *Photosynthesis in silico: Understanding Complexity from Molecules to Ecosystems* ed A Laisk and L NedbalGovindjee (Berlin: Springer) pp 231–46

Bernacchi C J, Singsaas E L, Pimentel C, Portis A R Jr and Long S P 2001 Improved temperature response functions for models of Rubisco-limited photosynthesis *Plant Cell Environ.* **24** 253–9

Bonan G B, Lawrence P J, Oleson K W, Levis S, Jung M, Reichstein M, Lawrence D M and Swenson S C 2011 Improving canopy processes in the community land model version 4 (CLM4) using global flux fields empirically inferred from FLUXNET data *J. Geophys. Res.* 116 G02014

Bonan G B, Oleson K W, Fisher R A, Lasslop G and Reichstein M 2012 Reconciling leaf physiological traits and canopy flux data: use of the TRY and FLUXNET databases in the community land model version 4 *J. Geophys. Res.* 117 G02026

Busch F A and Sage R F 2016 The sensitivity of photosynthesis to O₂ and CO₂ concentration identifies strong Rubisco control above the thermal optimum *New Phytol.* **213** 1036–51



- Busch F A, Sage R F and Farquhar G D 2018 Plants increase CO₂ uptake by assimilating nitrogen via the photorespiratory pathway *Nat. Plants* 4 46–54
- Ciais P et al 2013 Carbon and other biogeochemical cycles Climate Change 2013: The Physical Science Basis. Contribution of Working Group 1 to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change Chapter 6: Carbon and Other Biogeochemical Cycles ed T F Stocker, G K Plattner, S K Tignor, J Allen, A Boschung, Y Nauels and V B Xia (Cambridge: Cambridge University Press) pp 465–570
- Collatz G J, Ball J T, Grivet C and Berry J A 1991 Physiological and environmental regulation of stomatal conductance, photosynthesis and transpiration: a model that includes a laminar boundary layer *Agric. Forest Meteorol.* 54 107–36
- Collatz G, Ribas-Carbo M and Berry J 1992 Coupled photosynthesis-stomatal conductance model for leaves of C₄ plants *Aust. J. Plant Physiol.* **19** 519–38
- Dahal K, Wang J, Martyn G D, Rahimy F and Vanlerberghe G C 2014 Mitochondrial alternative oxidase maintains respiration and preserves photosynthetic capacity during moderate drought in *Nicotiana tabacum Plant Physiol.* 166 1560–74
- Ellsworth D S, Crous K Y, Lambers H and Cooke J 2015 Phosphorus recycling in photorespiration maintains high photosynthetic capacity in woody species *Plant Cell Environ*. 38 1142–56
- Farquhar G, Caemmerer Von S and Berry J 1980 A biochemical model of photosynthetic CO₂ assimilation in leaves of C₃ species *Planta* 149 78–90
- Gu L, Pallardy S G, Tu K, Law B E and Wullschleger S D 2010 Reliable estimation of biochemical parameters from C₃ leaf photosynthesis-intercellular carbon dioxide response curves Plant Cell Environ. 33 1852–74
- Harley P C and Sharkey T 1991 An improved model of C₃ photosynthesis at high CO₂—reversed O₂ sensitivity explained by lack of glycerate reentry into the chloroplast *Photosynth. Res.* 27 169–78
- Kattge J and Knorr W 2007 Temperature acclimation in a biochemical model of photosynthesis: a reanalysis of data from 36 species *Plant Cell Environ*. 30 1176–90
- Kattge J et al 2011 TRY—a global database of plant traits Glob. Change Biol. 17 2905–35
- Kattge J, Knorr W, Raddatz T and Wirth C 2009 Quantifying photosynthetic capacity and its relationship to leaf nitrogen content for global-scale terrestrial biosphere models Glob. Change Biol. 15 976–91
- Lawlor D W 2002 Limitation to photosynthesis in water-stressed leaves: stomata vs. metabolism and the role of ATP Ann. Bot. 89 871–85
- Le Quéré C et al 2016 Global carbon budget 2016 Earth Syst. Sci. Data 8 605–49
- Lombardozzi D L, Sparks J P, Bonan G and Levis S 2012 Ozone exposure causes a decoupling of conductance and photosynthesis: implications for the Ball-Berry stomatal conductance model *Oecologia* 3 651–9
- Lombardozzi D L, Bonan G B, Smith N G, Dukes J S and Fisher R A 2015 Temperature acclimation of photosynthesis and respiration: a key uncertainty in the carbon cycle-climate feedback *Geophys. Res. Lett.* 42 8624–31
- Lovenduski N S and Bonan G B 2017 Reducing uncertainty in projections of terrestrial carbon uptake *Environ. Res. Lett.* 12 0440208
- EL Maayar M, Ramankutty N and Kucharik C J 2006 Modeling global and regional net primary production under elevated atmospheric $\rm CO_2$: on a potential source of uncertainty *Earth Interact.* 10 1–20

- Meehl G A *et al* 2012 Climate system response to external forcings and climate change projections in CCSM4 *J. Clim.* **25** 3661–83
- Mercado L M, Huntingford C, Gash J H C, Cox P M and Jogireddy V 2007 Improving the representation of radiation interception and photosynthesis for climate model applications *Tellus B* 59 553–65
- Oleson K et al 2013 Technical description of version 4.5 of the community land model (CLM) NCAR Technical Note (Boulder, CO: National Center for Atmospheric Research) p 434
- Quebbeman J A and Ramirez J A 2016 Optimal allocation of leaf-level nitrogen: implications for covariation of $V_{\rm cmax}$ and $J_{\rm max}$ and photosynthetic downregulation *J. Geophys. Res. Biogeo.* 121 2464–75
- Reich P B, Wright I J and Lusk C H 2007 Predicting leaf physiology from simple plant and climate attributes: a global GLOPNET analysis *Ecol. Appl.* 17 1982–8
- Rogers A 2014 The use and misuse of $V_{c,max}$ in Earth system models *Photosynth. Res.* 119 15–29
- Rogers A *et al* 2017a A roadmap for improving the representation of photosynthesis in Earth system models *New Phytol.* **213** 22–42
- Rogers A, Serbin S P, Ely K S, Sloan V L and Wullschleger S D 2017b Terrestrial biosphere models underestimate photosynthetic capacity and ${\rm CO_2}$ assimilation in the Arctic New Phytol. 26 705
- Sage R F, Sharkey T D and Seemann J R 1989 Acclimation of photosynthesis to elevated CO₂ in five C₃ species *Plant Physiol*. 89 590–6
- Sage R F, Sharkey T D and Pearcy R W 1990 The effect of leaf nitrogen and temperature on the CO₂ response of photosynthesis in the C₃ dicot *Chenopodium album* L *Aust. J. Plant Physiol.* 17 135–48
- Sellers P J, Randall D A, Collatz G J, Berry J A, Field C B, Dazlich D A, Zhang C, Collelo G D and Bounoua L 1996 A revised land surface parameterization (SiB2) for atmospheric GCMs Part 1. Model formulation J. Clim. 9 676–705
- Sharkey T D 2016 What gas exchange data can tell us about photosynthesis *Plant Cell Environ.* **39** 1161–3
- Vassey T L and Sharkey T D 1989 Mild water-stress of phaseolus-vulgaris plants leads to reduced starch synthesis and extractable sucrose phosphate synthase activity *Plant Physiol*. **89** 1066–70
- von Caemmerer S 2000 Biochemical Models of Leaf Photosynthesis (Collingwood: CSIRO Publishing) pp 1–176
- Walker A P, Hanson P J and De Kauwe M G 2014 Comprehensive ecosystem model-data synthesis using multiple data sets at two temperate forest free-air $\rm CO_2$ enrichment experiments: model performance at ambient $\rm CO_2$ concentration *J. Geophys. Res. Biogeo.* 119 937–64
- Wang H, Prentice I C, Keenan T F, Davis T W, Wright I J, Cornwell W K, Evans B J and Peng C 2017 Towards a universal model for carbon dioxide uptake by plants *Nat. Plants* 3 734–41
- Watson-Lazowski A, Lin Y, Miglietta F, Edwards R J, Chapman M A and Taylor G 2016 Plant adaptation or acclimation to rising CO₂? Insight from first multigenerational RNA-Seq transcriptome *Glob. Change Biol.* 22 3760–73
- Wullschleger S 1993 Biochemical limitations to carbon assimilation in C₃ plants—a retrospective analysis of the A/C_i curves from 109 species *J. Exp. Bot.* 44 907–20
- Xu C, Fisher R, Wullschleger S D, Wilson C J, Cai M and McDowell N G 2012 Toward a mechanistic modeling of nitrogen limitation on vegetation dynamics PLoS ONE 7 e37914
- Yang J T, Preiser A L, Li Z, Weise S E and Sharkey T D 2016 Triose phosphate use limitation of photosynthesis: short-term and long-term effects *Planta* 243 687–98